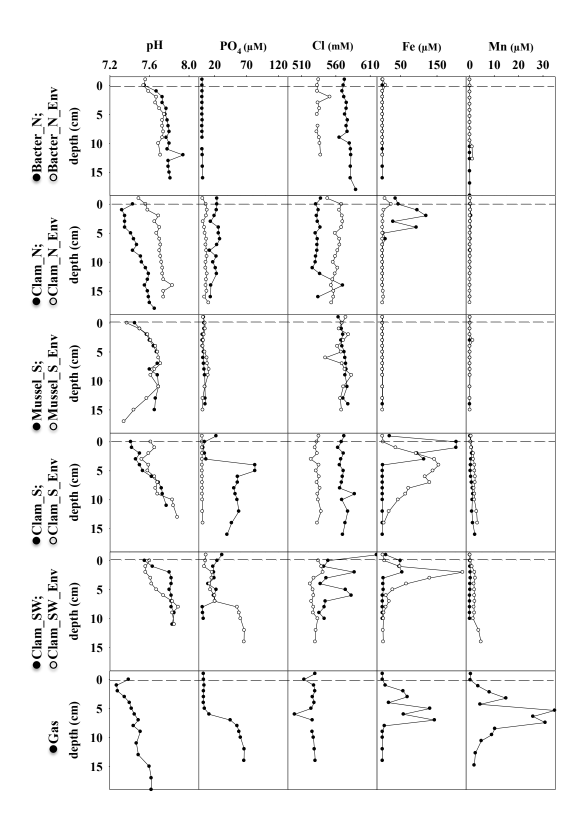
## **Supplement text 1:**

## Porewater geochemistry

Immediately after recovery and transfer of the push cores to *in situ* temperature of 4 °C, pH of every centimetre sediment depth was determined with punch-in electrodes on undisturbed sediment cores. Porewater analyses of phosphate and iron (Fe<sup>2+</sup>) were carried out on board. For the analyses of dissolved iron (Fe<sup>2+</sup>) porewater subsamples of 1 ml were immediately complexed with 50  $\mu$ l of "Ferrospectral" and determined photometrically. A photometric procedure was used to determine the concentration of phosphate in extracted porewater subsamples. Aliquots of the porewater were diluted 1:10 and acidified with HNO<sub>3 (suprapure)</sub> for the determination of manganese concentration by Inductively Coupled Plasma - Atomic Emission Spectrometry (ICP-AES) and Atomic absorption Spectroscopy (AAS) in the home laboratory.

Elevated alkalinity values were detected at all sites where also enhanced AOM activity was measured, indicating potential variation in the seepage between sites (Fig. 3). At these sites alkalinity increased with depth to reach maximum values of 47 and 49 mM in the deepest investigated horizons at the Mussel\_S and Bacter\_N sites, respectively (Fig.3). Highest alkalinity flux of 0.4 mmol m<sup>-2</sup> d<sup>-1</sup> was detected at the Mussel\_S and the Bacter\_N sites. Near background (2.5 mM) values were detected at the Clam\_S\_Env, Clam\_SW, Clam\_SW\_Env. The iron profiles matched well the sulphide depth pattern, with elevated concentrations detected at the clam habitats where free sulphide was absent from the topmost surface layers (Supplement Fig. 1).



**Supplement Fig. 1** Geochemical depth profiles of pH, PO<sub>4</sub>, Cl, Fe and Mn at all investigated sites at REGAB. Closed symbols denote measurements taken within the patches/bacterial mat, and open symbols denote measurements taken at the respective bare sediments.

**Supplement Table 1** Overview of the samples and measurements acquired at REGAB during M76/3b, with their PANGAEA reference numbers. All data has been deposited and is available online in the PANGAEA database (www.pangaea.de).

Location	Sampling site	Measurement; Sample	•
		DNA	M76/3b_310_PUC13; M76/3b_310_PUC27
	Bacter N	Porewater	M76/3b_310_PUC28; M76/3b_310_PUC32
	(Bacterial mat)	pH	M76/3b_310_PUC8
		AOM; SR	M76/3b_310_PUC27; M76/3b_310_PUC13; M76/3b_310_PUC12
		MICP	M76/3b_312_MICP1
		DNA	M76/3b_312_PUC7
	Bacter_N_Env	Porewater	M76/3b_312_PUC15
	(Outside bacterial mat)	pН	M76/3b_312_PUC34
		AOM; SR	M76/3b_312_PUC22; M76/3b_312_PUC23; M76/3b_312_PUC7;
N REGAB		DNA	M76/3b_323_PUC14
	CI N	Porewater	M76/3b 323 PUC15
	Clam_N	pН	M76/3b 323 PUC30
	(Clam patch)	AOM; SR	M76/3b 323 PUC28; M76/3b 323 PUC31; M76/3b 323 PUC14; M76/3b 323 PUC12
		CHAM	M76/3b 325 CHAM1
		DNA	M76/3b 332 PUC29
		Porewater	M76/3b 332 PUC31
	Clam_N_Env	pH	M76/3b 332 PUC20
	(Outside clam patch)	AOM; SR	M76/3b 332 PUC23; M76/3b 332 PUC29; M76/3b 332 PUC34
		MICP	M76/3b 335 MICP1
		DNA	M76/3b_344_PUC23
	Mussel S	Porewater	M76/3b_344_PUC30
	(Mussel patch)	pH	M76/3b_344_PUC29
		AOM; SR	M76/3b_344_PUC23; M76/3b_344_PUC28; M76/3b_344_PUC15
		CHAM	M76/3b_364_CHAM1
	Mussel S Env	DNA	M76/3b_361_PUC36
		Porewater	M76/3b_361_PUC14
	(Outside mussel patch)	pН	M76/3b_361_PUC13
	(Outside masser paten)	AOM; SR	M76/3b_361_PUC36; M76/3b_361_PUC15
		CHAM	M76/3b_364_CHAM2
S REGAB		DNA	M76/3b_361_PUC24
	Clam S	Porewater	M76/3b_361_PUC31
	(Clam patch)	pН	M76/3b_355_PUC9
		AOM; SR	M76/3b_361_PUC10; M76/3b_361_PUC21; M76/3b_361_PUC24
		CHAM	M76/3b_355_CHAM1
		DNA	M76/3b_355_PUC29
	Clam_S_Env (Outside clam patch)	Porewater	M76/3b 355 PUC35
		pН	M76/3b 355 PUC20
		AOM; SR	M76/3b 355 PUC11; M76/3b 355 PUC7; M76/3b 355 PUC29
		MICP	M76/3b 361 MICP1; M76/3b 361 MICP2
		CHAM	M76/3b 355 CHAM2
	Gas (Gas bubble)	DNA	M76/3b 364 PUC7
		Porewater	M76/3b 364 PUC21
REGAB		pH	M76/3b 364 PUC28
		AOM; SR	M76/3b 364 PUC7; M76/3b 364 PUC29; M76/3b 364 PUC9
	Clam_SW (Clam patch)	DNA	M76/3b 379 PUC28
SW REGAB		Porewater	M76/3b 379 PUC29
		pH	M76/3b 379 PUC34
		AOM; SR	M76/3b 379 PUC28; M76/3b 379 PUC14; M76/3b 379 PUC15
		CHAM	M76/3b_379_CHAM1
		DNA	M76/3b_379_PUC10
	CI CIVI E	Porewater	M76/3b_379_PUC9
	Clam_SW_Env	pH	M76/3b_379_PUC5
	(Outside clam patch)	AOM; SR	M76/3b_379_PUC10; M76/3b_379_PUC13
		MICP	M76/3b_385_MICP1; M76/3b_385_MICP2
		CHAM	M76/3b_379_CHAM2

**Supplement Table 2** Percentage of shared OTUs between all sites investigated at REGAB. Prior to this analysis, the depth samples within individual sites were merged.

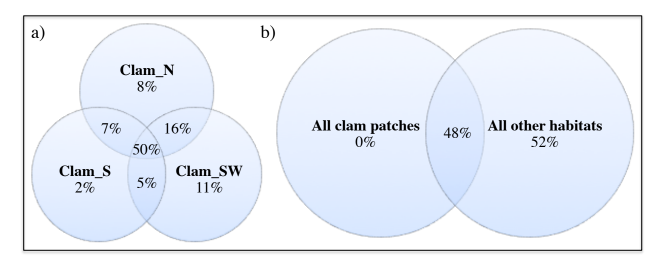
	Bacter_N	Bacter_N_Env	Clam_N	Clam_N_Env	Mussel_S	Mussel_S_Env	Clam_S_Env	Clam_S	Gas	Clam_SW
Bacter_N_Env	79									
Clam_N	83	73								
Clam_N_Env	84	77	82							
Mussel_S	69	65	67	71						
Mussel_S_Env	58	56	61	57	64					
Clam_S_Env	64	60	62	65	61	55				
Clam_S	68	64	69	68	66	57	67			
Gas	69	63	67	65	65	56	61	65		
Clam_SW	80	71	72	78	70	54	64	66	70	
Clam_SW_Env	73	65	69	72	68	53	60	63	65	79

**Supplement Table 3** Comparison of the shared OTUs (given as percentage) among all depth samples, between bare sediment sites and clam populated sites. An OTU was regarded as shared only if it was present in all samples (0-10 cm or 0-5 cm). The percentage of shared OTUs was calculated as the fraction of the total OTUs at the individual habitat.

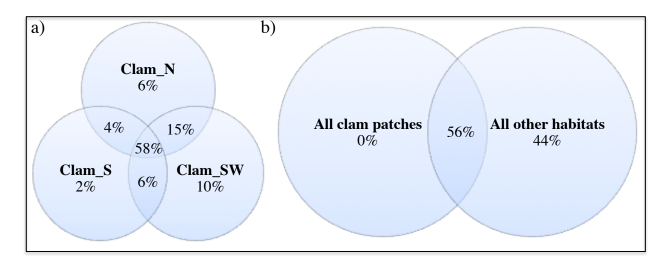
	Clam_N	Clam_S	SW_Clam	Clam_N	Clam_S	SW_Clam
	(0 - 10 cm)	(0 - 10  cm)	(0 - 10  cm)	(0 - 5 cm)	(0-5  cm)	(0 - 5 cm)
Inside	36	30	22	47	40	47
Outside	34	30	36	42	44	44

**Supplement Table 4** Distance-based test for homogeneity of multivariate dispersions. Table comprises average distances to the centroids, calculated based on Jaccard and Bray-Curtis dissimilarity indices. The higher the value of the average distance to the centroid, the higher the dispersion (variance) within the respective group. The test was performed incorporating only the surface samples (0-5 cm), or samples from all depths (0-10 cm).

	Clam_N	Clam_N_Env	Clam_S	Clam_S_Env	Clam_SW	Clam_SW_Env
<b>0 - 5 cm</b> (Jaccard)	0.3	0.3	0.4	0.3	0.3	0.3
0 - 5 cm (Bray-Curtis)	0.2	0.2	0.3	0.2	0.2	0.2
<b>0 - 10 cm</b> (Jaccard)	0.4	0.4	0.4	0.4	0.4	0.4
0 - 10 cm (Bray-Curtis)	0.2	0.2	0.3	0.3	0.3	0.3



**Supplement Fig. 2** OTU partitioning analysis taking into account all sediment depth samples (0 - 10 cm). 50% of the total OTUs were found to be shared by all three clam patches (a). The clam patches had no unique OTUs relative to the other investigated habitats at REGAB (b).



**Supplement Fig. 3** OTU partitioning analysis taking into account the topmost 5 cm sediment depth samples. 58% of the total OTUs were found to be shared by all three clam patches (a). The clam patches had no unique OTUs relative to the other investigated habitats at REGAB (b).